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- (54) RADIATION-CURABLE INK COMPOSITION STRAHLUNGSHÄRTBARE TINTENZUSAMMENSETZUNG
- COMPOSITION D'ENCRE DURCISSABI E PAR EXPOSITION A UN RAYONNEMENT (84) Designated Contracting States:
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The file contains technical information submitted after the application was filed and not included in this specification

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Description

[0001] The invention relates to a radiation curable inkcomposition, a method for preparing a coated and colored optical glass fiber, a coated and colored optical glass fiber, and a ribbon-assembly comprising such coated and colored optical riass fibers.

[0002] Optical glass fibers are generally costed with two superposed radiation-cured coatings, which together form a primary costling. The coating which is in direct constant with the glass is called the interprintery coating and the overtaying coatings is called the such primary coating and the overtaying coatings is called the soundary coating. The interprimary may also be called the primary coating, then, the outer arimary coating is called the soundary coating. Sold definitions are used interchangeable.

10 [0003] The inner primary coating is usually a relatively soft coating providing environmental protection to the glass fiber and resistance, inter alia, to the well-known phenomenon of microbending. Microbending in the coated fiber can lead to attenuation of the signal transmission capability of the coated fiber and is therefore undestraine. The outer primary coating(s), which is on the exposed surface of the coated fiber, is typically a featively harder coating designed to provide a diselect explanation.

16 [0004] In telecommunications applications of optical fibers, multiple individual strands of coated fiber must be eackaged into larger structures such as ribbons and coables to maximize efficiency. However, after ribboning and cabling of fiber, the individual strands of fiber must be readily distinguishable from each other so they can be accurately identified uring, for example, installation and repair. Outsile geometry and/or color coding can be used to distinguish and identify individual fibers in a complex cable. Although several methods can be used to color code fiber, color coding can be done avoirable appounds with either a thin ink layer (< 10 microns) which is placed over the coated fiber before ribboning and cabling or bu sinc a colored outer finance ordinary coatino.</p>

1900g1 Tape-like optical their ribbons are prepared by embedding a plurality (e.g., 4 to 12) of individual color coded fibers in a supporting matrix material which, like the inner and outer primary countings, is also anolation-curable to maximize production speed. The matrix material can enge-bond of the glass fibers to gather. Ourse of the matrix material county of potential gas after the fibers have been color coded by ink. Hencer, in a ribbon design, the his layer residue between the fibers matrix material and the fibers' outer primary costing. This means that the ink layer's interfacial characteristics (e.g., surface energy, echesion) must be carefully controlled to function properly with both matrix material and outer primary costing in the ribbon structure. In a structure, the ability of a curred matrix material to be suitably stripped off the ink layer foreact-out) is an important technical some structure.

with use of solvents is also known.

[0006] Octofal fiber color coding can be based on up to 12 or more colors. Although optical fiber inks were originally solvent-based or thermosetting inks, in more recent times, radiation-curable inks have been used to increase the speed of the inking process. In those ink compositions, plyment is dispersed in a radiation-curable carrier or base composition.

[0007] As the demand for coated optical glass fibers has increased, manufacturers must respond by adding more fiber drawing production fines and by strengting to increase the finear line speeds of the existing fiber drawing production fines. In the stater case, one factor within will determine the upper limit for the line speed will be the curing rate charges.

teristics of the rediation-curable ink composition, for a given radiation source and intensity.

[DOO8] If the fine speed is increased to the extent that cure rate there requirements of the radiation-curable ink composition are not provided, the radiation-curable ink composition will not have received a sufficient amount of radiation to cause complets cure, or cross-linking, of the radiation-curable ink composition. The production interer line speed is generally inversely related to the amount of radiation explosure to the radiation-curable ink composition during the production process will necessarily decrease for a given radiation source, incomplete cure of the radiation-curable ink composition is undefined to the radiation and the second control or process will necessarily decrease for a given radiation source, incomplete cure of the radiation-curable ink composition is undefined to the radiation and the second control or control or the radiation source.

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[0009] In general, reduction-curable ink coating compositions can cure at a significantly slower rate than radiation-curable outer primary coating compositions, it is believed that the pigments present in ink compositions contribute to the slower cure speed of link coatings. Thus, there is a need for improving the cure speed of the ink coating.

[0010] While the ink composition must have a very fast cure speed to ensure complete cure of the link coating on the high speed drawing units, the increase in our speed should not come at the expense of other important properties of the ink coating, such as providing suitable break-out performance. Break-out performance is the ability of the cured ink coating to separate from the markix naterial without separating the ink layer from the outer primary coating to providing mid-span access to the individual coated optical gises there contained within the ribbon reasembly. Mid-span access is access to the coated optical gises fibers at a portion of the ribbon assembly between the ends of the ribbon assembly. Therefore, there is a need for a redistriou cruzible ink composition that exhibits adaptable adhesion properties to provide

an adhesion between the outer primary coating and the ink coating that is greater than the adhesion between the ink coating and the matrix material to provide mid-span access.

[0011] In addition, ink compositions should not contain ingredients that can ingrate to the surface of the optical glass fiber and cause corresion. The ink composition should ask on cloratian ingradients which can cause instability in the protective coatings or matrix material. Ink coatings for optical glass fibers should be color fast for decades, not exuse attenuation of the signal transmission, be impervious to cabling gels and chemicals, and allow sufficient light penetration for fiber core slignment.

[0012] Analogous requirements exist for colored outer primary coatings. In particular, these coetings require high cure speed desire the presence of prigments, good teselved not properlies, opport demonstrate the inner primary coating, and the coeting should contain no ingredients that may cause attenuation of light through the glass fiber after prolonged exposure of the ribbon to moist. heat, etc.

[0013] From the above, it is clear that optical glass fiber technology places many unique demands on radiation-curable colored compositions which more conventional technologies, such as printing links, do not.

[0014] U.S. patent No. 4,629,265 discloses a method for making an ink coating on a coated optical glass fiber in which a UV curable ink is applied to a coated optical glass fiber. The Ink coating is applied in a method that preserves the concentricity of the optical glass fibers. The preferred inks are pigmented semi-lopaque UV curable polymetric lists. However, the link compositions disclosed in this patent do not have a sufficiently fast enough cure speed to be used on modern high speed outside uses fiber drawing and coating tower.

(0015) Published Japanese Patent Application No. H1-152405 discloses a radiation-curable ink composition containing an organic polylioxane compound. The polysiloxane compound provides the Ink coating with the ability to separate more easily from the matrix material in a riboon assembly.

[0016] Published Japanese Patent Application No. 64-22976 discloses radiation-curable ink compositions containing specific radiation-curable oligomers. The ink composition provides an Ink locating having adhesion to the outer primary coating which is separable from the matrix meterial in a ribbon assembly.

29 [0017] Patent application EP-A-B1099 describes the use of a release agent such as a allicon oil or a fluororesin between the bundling layer and the coloring layer. In particular, when adustantial manurat of silicone resine are used, incompatibility in the liquid and resultant imperfections in the cured matrix composition may result, which causes attenuation of line.

(0018) Usually link compositions must be cursed in an inert atmosphere, i.e. in the ebsence of oxygen. Providing inert strongshrees to optical glass filter drawingtowers is expensive. Thus, an antidisnor-curstle ink composition which exhibits a high curse speed in the presence of some oxygen like in an air atmosphere would provide significant advantages over ink compositions that must be used in an inest atmosphere.

[0019] An objective of the present invention is to provide ribbon assemblies having suitable break-out performance, without requiring the use of substantial amounts of a silicone agent in the ink composition.

59 [0020] Another objective of the present invention is to provide radiation-curable ink compositions which are adapted to provide cured ink coatings having greater adhesion to the outer primary coating than to a matrix material, to provide the ribbon assembly with the capability of break-out of the individual coated optical glass fibers.

[0021] A further objective of the present invention is to provide high speed radiation-curable ink compositions which provide enhanced cure speed in the presence of air.

40 [0022] Another objective of the present invention is to provide an ink composition which exhibits enhanced adhesion to the outer primary coating when suitably cured.

[0023] Other objectives of the prasent invention are to provide an optical glass fiber coated with the lnk composition and a ribbon assembly containing at least one of the coated optical glass fibers.

[0024] Provided by the present invention is a radiation-curable ink coating composition comprising

- a) an oligomer
- b) a reactive diluent
- c) a homolytic photoinitiator
- d) a pigment, and
 - e) additives

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wherein the composition comprises more than about 1 wt.% homolytic photoinitiator in combination with a benzophenone-type compound, as defined in claims and dependent claims 2-0. Also provided by the present invention is a method for curing the radiation curable ink composition as defined in claim 10, a coated and colored optical glass fiber as defined in claims 11-14 and a ribbon assembly as defined in claim 15. The novel ribbon assembly his those but capability of providing break-vot or clore coded optical glass fibers from the ribbon assembly, the ribbon assembly homolying a purally of coated optical glass fibers then the ribbon assembly, the ribbon assembly comprising a purally of coated optical glass fibers hering an ink or colored secondary coating on at least one of the coated optical glass fibers and a matrix material binding the plurality or obserted optical prices together.

[0025] This composition if suitably formulated, using the constituents a-e, can also be used as a colored outer primary coating.

[0026] Unexpectedly, this use of a benzophenone-type compound - if suitably cured - results in a system having a satisfactory adhesion to an outer primary coating and good break-out from the matrix material.

5 [0027] In DE:A-4007519, it is described to use photoinitiators in the curing of UV curable ink compositions on optical fibers. However, this reference does not teach to use the combination of a substantial amount of homolytic types of photoinitiators in corribination with substantial amounts of benzohenone.

[0028] The radiation-curable ink compositions comprise two functional components: a pigmant system which functional components: a pigmant system which functions to allow the pigment system into a coherent, or independent of the pigment system into a coherent, or independently film after radiation-cure. Typically, a carrier system will be mixed with a pigment system to form the radiation-cure in the pigmant system.

[0029] Ink coeffings are usually about 3 to about 10 microns thick, and should be concentric to prevent attenuation of the signal transmission. The ink coeting also generally has a Γ_{ij} of at least about 50°C. replaced transmission. The ink coeting also generally are 10-30 μ m thick, and are described e.g. in FB-8-478445.

6 [0031] Radiation-curable carrier systems which are suitable for forming the present ink composition contain one or more radiation-curable oligomers and monomers having at lease one fundancing proup capable of polymeratation when exposed to actinic radiation. Suitable radiation-curable oligomers or monomers are now well known and within the skill of the art.

[0032] The coated optical glass fibes each have an ink coating for identifying the individual fibers. The level of adhesion between the link coating and the matrix material has been adjusted to be less than the level of adhesion between the ink coating and the secondary coating on the optical glass fiber, which provides the functional capability of providing break-out of the individual coated optical glass fibers.

[0033] Ink compositions according to the present invention may show a certain percent reacted acrylate unsaturation ("Sh RAU") by selecting monomers, oligomers and a photolitation vehicle provides ShRAU", when exposed to about 0.125 Jichn² UV redation, such that the level of adhesion between the ink coating and the matrix material is less than the level of adhesion between the ink coating and the matrix material is less than the level of adhesion between the ink coating and the outer primary coating, Preferably, the everage ShRAU is at least 75%, more preferably at least about 80%, when cured in the presence of an oxygen cortaining gaseous atmosphere, Rice air. Preferably, the everage ShRAU is at least 75% more preferably at least about 80%, and most preferably at least about 80%, when cured in an Inent (e.g. thropen) atmosphere. The everage ShRAU is at least 50% and most preferably at least 40% at least 50% at l

[0034] Preferably, all three carbon containing compounds "Ar" in the homolytic photoinitator according to formula [1] in claim 6 have a molecular veright and molecular structure that is suitable for forming the above free-readicals. For example, the carbon containing compounds can have from about 5 to about 15 carbon atoms. Suitable examples of the carbon containing compounds include a furair enform or a benzene into.

[0035] One or more of the hydrogens present on the aromatic ring can be substituted, for example, with alkyl and/or alkoxy groups. Examples of suitable alkyl substituents include methylene, ethylene, burylene and propylene groups. Examples of suitable alkoxy substituents include methoxylete, ethoxylete, butoxylete and propoxylete groups.

40 [0036] A preferred cure speed enhancing photoinitiator includes one represented by the following formula (2):

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wherein Ar is a phenyl group, optionally substituted with one to three alkyl groups having 1-4 carbon atoms.

[0037] Suitable amounts of the photoinistors economing to formula (1) in claim is are from about 1 to about 25 weights, norterably from about 1,5 to about 20 weights, and most preferably from about 4 to about 15 weights, based on the total weight of the ink composition in general, if other photoinistors are present in the ink composition in combination with the ink cure speed enhancing photoinistor, the ink cure speed enhancing photoinistor can be used in lesser amounts to provide the desired cure speed.

[0038] Ink coatings are usually about 3 to about 10 microns thick, and should be concentric to prevent attenuation of the signal transmission. The ink coating also generally has a Tg of at least about 50°C, more preferably at least about 50°C.

[0039] Commonly, the radiation-curable functionality used is ethylenic unsaturation, which can be polymerized through radical polymerization or cationic polymerization

[0040] A suitable radiation-cumble ink composition essentially consists of from about 1 to about 80 weight % of at least one radiation-cumble oligomer include from about 20 to about 70 % by weight, based on the total weight of the link composition.

[0041] A mixture of mono-, di-, toi-, tetra-, and higher functionalized oligomers can be used to achieve the desired balance of properties, wherein the functionalization refers to the number of radiation-curable functional groups present in the oligomer.

[0042] The oligomers usually comprise a carbon-containing backbone structure to which the radiation-curable functional group(s) are bound. Examples of suitable arabon-containing backbone can be selected to provide fine, polyseties, polysmibes, and polycarbonates. The size of the carbon-containing backbone can be selected to provide the desired molecular weight. The number average molecular weight of the oligomer's usually between about 500 to about 10,000, preferably between about 500 to about 7,000, and most preferably between about 500 to about 7,000, and most preferably between about 500 to about 7,000, and most preferably between about 500 to about 7,000, and most preferably between about 500 to about 7,000, and most preferably between about 500 to about 7,000 and most preferably between about 500 about 7,000 about 5,000 about 7,000 about 5,000 about 7,000 about 5,000 about 5,000 about 7,000 about 7,000 about 5,000 about 7,000 about 5,000 about 7,000 about 5,000 about 7,000 abou

[0043] For example, the carbon-containing backbone of the oligomer can comprise aromatic groups and ring-opened epoxy groups or alkoxy groups. The oligomer can be represented by, for example:

B-Ar-B

or

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R-I -Ar-I -R

where R is a radiation-curable functional group,

Ar is an aromatic group containing molety, and

25 L is a linking group.

[0044] Examples of suitable linking groups include alkoxy or ring opened epoxy such as ethoxy, propoxy, butoxy, and repeat units thereof. L. can also be a urethane or urea linking group.

[0445] The aromatic groups can be, for example, derived from bisphenol units, such as bisphenol A. A preferred of logomer is a diplyridly field redivative of bisphenol A to which excityful functional groups have been bound. A commercially example of such an oilgomer is Cberryl 3700 (UCB) or CN-120 (Sintomer), the latter having a molecular weight of about 1500.2 and when CM or observed to the control of the c

[0046] Another example of a preferred oligomer is a trifunctional polyether or polyester having a molecular weight of about 500 to about 5000. A preferred example of a trifunctional oligomer is the commercially available polyurenane triacrylate Ebecn/1 264, which has a molecular weight of about 2000 and when cured a Tig of about 42°C.

- 5 (0047) The radiation-curable carrier systems may also contain a reactive diluent (b) which is used to adjust the viscosity. The reactive diluent can be a low viscosity monomer containing having at least one functional group capable of polymerization when exposed to action cradiation. This functional group may be of the same nature as that used in the radiation-curable monomer or oligomer. Proferably, the functional group present in the reactive diluent is capable of copolymentizing with the radiation-curable monomer or oligomer.
- 40 [0048] A suitable madiation-curable ink composition essentially consists of from about 1 to about 80 weight % of at least one radiation-curable diluent. Preferred amounts of the radiation-curable diluent include from about 5 to about 80 % by weight, more preferably from about 10 to about 80 % by weight, based on the total weight of the ink composition. [0049] Preferably, compound b) is a reactive diluent or a mixture thereof. Generally, each reactive diluent has a molecular weight of the stant about 550 may alwoish of less than about 550 may alwossibly of less than about 550 may alwossibly of less than about 550 may alwossibly disease.
- 45 [0050] For example, the reactive diluent can be a monomer or mixture of monomers having an acrylate or vinyl ether functionality and a C4-C20 alkyl or polyether moiety.
 - [0051] Particular examples of such reactive diluents include: hexylecrylate, 2-ethylhexylecrylate, isobornylacrylate, decylacrylate, isobornylacrylate, isoprolacram, N-vinylpytrolidone, and the like.
 - [0052] Another type of reactive diluent that can be used is a compound having an aromatic group. Particular examples of reactive diluents having an aromatic group include: ethyleneglycolphenylether-acrylate, polyethyleneglycolphennylether-acrylate, polypropyleneglycolphenylether-acrylate, and alkylsubsitiuted phenyl derivatives of the above monomers, such as polyethyleneglycolnonylphenyletheracrylate.
 - [0053] The reactive diluent can also comprise a diluent having two or more functional groups capable of polymerization.

 Particular examples of such monomers include:

C2-C18 hydrocarbon-diololiacrylates,

C₄-C₁₈ hydrocarbondivinylethers,

- C3-C18 hydrocarbon triacrylates, and the polyether analogues thereof, and the like, such as
- 1,6-hexanedioldlacrylate,
- trimethylolpropanetri-acrylate,
- hexanedioldivinylether,
- triethylene-glycoldiacrylate, pentaerythritol-triacrylate.
 - ethoxylated bisphenol-A diacrylate, and
 - tripropyleneglycol diacrylate.
- 10 [0054] Preferably, alkoxylated aliphatic polyacrylates are used, such as ethoxylated hexanedioldiscrylate, propoxylated glyceryl triacrylate or propoxylated trimethylolpropanetriacrylate.
 - [0055] At least one of the photoinitiators of used in the link coating composition of the present invention is a homolytic fragmentation photoinitiator which operates by intramolecular bond cleavage,
 - [0056] A suitable radiation-curable ink composition essentially consists of from about 1 to about 20 weight% of a homolytic photoinilator, Preferred amounts of the homolytic photoinilator are from greater than 3 to about 15% by weight, more preferably from about 4 to about 12% by weight.
 - [0057] As a pigment is present in the composition according to the Invention, it is preferred (also for colored outer primary coating compositions) to use as shotchritistor an eavy phosphine existe photchritistor, such as defined in claims 6 and 7, more specifically a benzoyl d'aryl phosphine oxide photoinitistor. Examples of suitable benzoyl d'aryl phosphine oxide photoinitiations include: 2.4.6-timetriy/benzoyldiphenry/hosphine oxide (Luclin TPO by BASF), and
- oxide pnotonitators include: 2,4,6-timethyloentoylidipherely-phosphine oxide (Lucirin TPO by BASF), and bis(2,6-dimethoxyloenzoyl)-2,4,4-trimethyloentyl-phosphine oxide (Ingacure 1700 by Clba Gelgy).
 [0058] For an optimum cure speed in the presence of bioment, it is advantageous to combine an acyl phosphine oxide
 - judooj indra opurnum cure speed in the presence or pigment, it is advantageous to combine an acyl phosphine oxide photoinitiator with one or more other photoinitiators, such as hydroxy-cyclohexylphenyl-ketone.

 [1059] Any inorganic and organic bigment (d) that is suitable for making radiation-curable ink compositions can be
- 25 used in the present invention. However, because white pigments usually do not absorb splintions and compositions can be used in the present invention. However, because white pigments usually do not absorb splintions amounts of the calabilities used to cure the ink composition, white ink compositions do not usually require the use of the ink cure speed enhancing photoinitiation according to the present invention. Therefore, he preferred bigments are pigments that absorb light of a visible wave length, i.e. any color except pure white. The use of the term "pigment" refers to both inorganic and organic pigments.
- 90 [0060] Preferably, the pigment used in the link coating composition of the present invention is an organic pigment. The pigment can be present in the link composition in an amount that provides coloration that is visible without magnification to facilitate identification of the individual colored optical classis fiber.
- [0061] Ribbon assemblies utifizing 12 or less coatled optical glass fibers require only 12 colors to adequately distinguish each of the coated optical fibers from one another. However, in larger ribbon assemblies, more than 12 colors may be sufficed to adequately distinguish the coated optical glass fibers from one another. Examples of twelve colors normally used for making ribbon assemblies include: black, white, yellow, blue, red, green, orange, brown, pink, aqua, violet, and
 - [0062] Preferably, the pigment has a mean particle size of not more than about 1 µm. The particle size of the commercial pigments can be lowered by milling if necessary.
- 40 [0063] A specific example of a suitable black pigment includes carbon black.

ments, quinacridone pigments, anthraquinone, and perylenes.

- [0064] A specific example of a suitable white pigment includes titanium dioxide.
- [0065] Specific examples of suitable yellow pigments include diarylide yellow and diazo based pigments.
- [0066] Specific examples of suitable blue pigments include that yield yield wand diazo dased pigments.
- anines.

 45 [0067] Specific examples of suitable red pigments include anthraquinone (red), napthole red, monoazo based pig-
 - [0068] Specific examples of sultable green pigments include phthalocyanine green and nitroso based pigments.
 - [0069] Specific examples of suitable orange pigments include monoazo and diazo based pigments, quinacridone pigments, anthraquinones and pervienes.
- 50 [0070] Specific examples of suitable violet pigments include quinacrinode violet, basic dye pigments and carbazole dioxazine based pigments.
 - [0071] Suitable aqua, brown, gray, and pink colors can easily be formulated by combining several pigments. One skilled in the art is able to form any color as desired by combining different pigments.
- [0072] The pigment can be present in the ink composition in an amount that provides coloration that is visible without magnification to facilitate Identification of the individual colored optical glass fiber. The amount of the pigment should not be a great as to significantly reduce the cure speed of the ink composition or result in other undesirable affects. Examples of suitable amounts of pigment have been found to be higher than about 1-25 weight %. Generally, the amount is loss that 25 w.fx. preferably less then about 10 weight %. Beach over orderably less than about 10 with %. Beach or the preferably less than about 10 weight %. Descriptions.

the total weight of the ink composition.

[0073] Preferred amounts of each pigment are from about 0.5 to about 10 % by weight, more preferably from about 0.5 to about 5 % by weight, in colored outer primary coatings, generally lower amounts of pigments are sufficient to achieve good coloring.

- 5 [0074] Other additives which can be used in the radiation-curable carrier system include, but are not limited to, lubricants, wetting agents, entiocidusts and stabilizes. The selection and use of such additives is within the skill of the art. [0075] Coated optical fibers are often used in ribbon assemblies. Because of the versatility of the presently invented in incoating composition, this composition is very well butted for use on contact optical glass bloers in ribbon assemblies. If necessary, a release agend can be added to the ink costfing to allow easy access to the individual Bitters by separafilling the matrix material from the ink costing, usually called break-out. However, preferably, no substantial amount of release agents include silicones, silicone caryitates, fluor-carbon oils or realism and the like, if present, the ink costing composition for ceating optical fibers used in ribbon assemblies comprises about 0.1 to about 20 w.%, more preferable bout 1.1 to about 1.0 which 1.0 w.%, of a suitable release agent is based on the total work of the time.
- composition.

 [1076] Useful additives e) of the inx coating are a small amount of lubricant to allow even better access to the individual fibers in separating the matrix material from the inx coating, usually called break-out. Suitable lubricants incluse silicones, rours carbon oils or resins and the like. Reactive release agents such as allicone acquites appear to have no further effect. Preferably, the inx coating composition for coating optical fibers used in ribbon assemblies comprises about 0.01 to about 5 wfx, more preferably about 0.1 to about 5 wfx fx, more preferably about 0.1 to about 5 wfx fx, more preferably about 0.1 to about 5 wfx fx fx.
 - composition.

 Preferably, a silicon oil is used. Another jubricant compound can be present to a lesser extent.

[0077] The link coating composition of the present invention comprises an effective amount of a benzophenone-type compound to achieve a better break-out of the markin material from the coated and colored optical files, it auticated addition-outed ink composition of this embodiment comprises more than about 2 weight % of a benzophenone-type

[0078] Preferred amounts of the benzopherone-type compound are more than about 5 % by weight, and more prefarably more than 3.5 % by weight, based on the batal weight of the ink composition. The amount generally will be less than about 10 % by weight as use of more than about 10 % by weight of the benzopherone-type compound creates no periodiar advantage, and higher amounts are not recommended from economical point of view. Preferably, the ink composition composition comprises less than 6 wt% of benzopherone-type compound.

[0079] Suitable benzopherone-type compounds are organic compounds with a malecular weight between 170 and sometic and which consists essentially of a substituted benzopherone group. Sultable substituents are allyl groups, acry groups, armatic groups, halogens, and the like. Suitable benzopherone-type compounds include benzopherone, chlorobenzopherone, methyl-o-benzoyl benzoate, 4-benzoyl-4-methyldiphenylsulphide, acrylated benzopherone, 4-phenylben-

5 zophenone, 3,3 dimethyt-4-methoxybenzophenone, 4,4-dimethylamino-benzophenone, and the like. Preferably, this benzophenone-type compound is not a ring-bonded benzophenone. Fling-bonded benzophenones such as thioxanthone or derivatives thereof, e.g. 2,4-diethyl- or 9-isopropyl-thioxanthone, appear to have little effect on the improvement of break-out properlies.

[0080] The ink composition according to the present invention preferably comprises none, or a minor amount of amine compound, which would act as a hydrogen abstraction co-initiator for the benzophenone type compound.

[0081] Generally, less than 0,5 wt.% and preferably less than 0,1 wt.% of amine type hydrogen abstraction compound would be present in the composition of the present invention. Nevertheless, morpholino-type sensitizers can be present in amounts up to 4 wt.% but preferably less than 3 wt.%.

[0082] Based on the above, a suitable radiation-curable ink composition can be formulated from a composition con-

from about 1 to about 20 weight % of at least one pigment that absorbs light of a visible wavelength; from about 1 to about 25 weight % of at least one photoinitiator; and

from about 55 to about 98% of a radiation-ourable carrier system containing at least radiation-ourable monomer or oligomer, based on the total weight of the ink composition.

[0083] A preferred ink composition can be formulated from a composition containing:

from about 1 to about 20 weight % of at least one pigment which absorbs light of a visible wavelength;

from about 1 to about 80 weight % of at least one radiation-curable oligomer,

from about 1 to about 80 weight % of at least one radiation-curable diluent monomer; and

from about 1 to about 20 weight % of at least one ink cure speed photoinitiator, based on the total weight % of the ink composition.

[0084] Preferred amounts of the ink cure speed enhancing photoinitiator are from about 1.5 to about 20 weight %, more preferably from about 4 to about 15 weight %.

[0085] Preferred amounts of the radiation-curable oligomer include from about 10 to about 70% by weight, more preferably, about 20 to about 60% by weight, based on the total weight of the ink composition.

- 5 [0086] Preferred amounts of the radiation-curable diluent monomer include from about 10 to about 70% by weight, more preferably, about 20 to about 60% by weight, based on the total weight of the ink composition.
 - [0087] Preferably, the ink composition contains other photoinitiators in an amount of about 1 to about 20% by weight, more preferably about 1 to about 10% by weight, based on the total weight of the ink composition.
 - [0088] Raddrion-cure of optical fiber inks can be executed with radiation sources which have characteristic emission outcuts which will vary depending on the radiation source. Examplary radiation sources can be obtained from, for example, Fusion UV Systems inc. or Iwasaki Denki Co. For example, so-called "D-lamps" and "H-lamps" can be used as a radiation source, wherein the D lamp is more rich in longer wavelength uttraviolet light emissions than the H lamp. A typical uttraviolet radiation source (e.g., medium pressure and tamp) used in Industrial production often has major emission lines near the 280, 300, 313, and 368 mm ranges, or more generally at about 250-270 mm, at about 290-320
- emission lines near the 280, 300, 313, and 386 m ranges, or more generally at about 260-270 mm, at about 290-320 nm, and at about 360-380 nm. In addition, there can be a line at about 390-410 nm. The characteristic bub o upput used in the present invention can be measured by conventional methods or obtained from the supplier of the radiation source. Preferably, the ink composition of the present invention is cured by irradiation with both an H and a D lamp.

[0089] The colored coating compositions of the present invention preferably are cured with sufficient energy, to have lost than 15% unreacted acrylate unsaturation. Suitable irradiation energy values are 0.05-2 J/cm² preferably 0,1-1.5 J/cm². More in particular, values below 0.5 J/cm² are particular preferred.

[0090] The present invention also provides an ink composition having enhanced adhesion to the outer primary coating when suitably cured. It has been found that surprisingly the adhesion of the ink coating to the outer primary coating can be approximated by how well a monomer present in the ink composition adsorbs in the outer primary coating, in general, the greater the adsorption of the monomer into the outer primary coating, the greater the adhesion of the dured ink coefficiant the fourth primary coating.

[0091] If the link coating separates from the outer primary coating during mid-spen access to the individual cotted glass filters, the identification of the individual optical glass filters can be difficult or even unstalentable. Using the present invertion, the achiesion of the link coating to the outer primary coating can be easily adjusted to provide an ink coating having greater adhesion to the outer primary coating than to the matrix metarial to provide a ribton seasonably that is capable of providing mid-span access to the individual optical glass fibers while avoiding undesirable delamination of the ink coating from the outer primary ocating uniter mid-span access.

[0092] Preferably, the monomer present in the link composition exhibits an Outer Primary Adsorption Index of at least about 5, preferably at least about 7, and most preferably at least about 7.

[0093] The adsorption index of the monomer can be determined without undue experimentation by first preparing 75 in micron thick diswaldown of the outer primary conting to be coated by the link composition. The drawdown is then suitably curved by excessure to redistion and then cut into 2 inch by 2 inch sample squares. The sample square is then submerged in the monomer present in the desired link coating for 50 seconds under atmospheric pressure and nown temperature. (25°C). The amount of weight gath in the sample expressed as a percentage of the original weight of the sample is used to determine the outer primary adsorption index as follows.

40 [0094] Outer Primary Adsorption Index;

- 1 = 1% weight gain
- 2 = .2% weight gain
- 3 = .3% weight gain
 - 4 = .4% weight gain
 - 5 = .5% weight gain
 - 6 = .6% weight gain
 - 7 = .7% weight gain
 - 8 = .8% weight gain
 - 9 = .9% welght gain
 - 10 = 1% weight gain

[0095] Thus, for an adsorption index of 5, the sample gained about .5% in weight after being submerged in the monomer for 30 seconds.

50095 The amount of the monomer can be selected to provide the destred level of adhesino between the outer private coeting and the link coating, in general, the greater the amount of a monomer having an adexprible index of at least so about 5, the greater the adhesion between the outer privary creding and the link coating. Suitable examples of such a monomer are from about 1 to about 20 wt. 8, and preferably from about 15 with coating.

of the ink composition.

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[0997] In producing a coated and colored optical fleer, a liquid coating composition is applied to a substrate and subsequently cured. The optical glass floers are generally coated with two superposed radiation-cured coatings, which together form a primary coating. The finer primary coating is in direct contact with the glass and the outer primary coating is adjacent to the surface of the inner primary coating. The fine coating composition according to the present invention can be formulated to provide an fine coating how a good adhesion to the outer primary coating.

[0098] It is also possible to use a colored outer primary osetting, which prefutes the necessity of using an extra ink inyer. The outer primary ceating should adhere well to the inner primary ceating, and the matrix methed is hould show good break-out properties with respect to the colored outer primary ceating, which can be achieved using the ink ceating of the present invention as the colored outer primary ceating.

[0099] The above embodiments can be used in combination to provide an improved radiation-curable ink composition which exhibits the combination of properties of:

- (i) when suitably cured provides the ribbon assembly with the functional capability of providing break-out without detaminating the ink or colored coating from the outer resp. inner primary coating;
- (ii) when suitably cured provides sufficient adhesion between the matrix material and the ink coating to resist deta-
 - (iii) exhibits an enhanced cure-speed in the presence of an oxygen containing atmosphere such that the colored coating can be formed at high speed without the need for providing an inert atmosphere.
- [0100] The ink coating composition can be applied to the coated optical glass and cured using any suitable method. An example of a suitable method is disclosed in U.S. patent No. 4,629,285, the complete disclosure of which is incorportated herein by reference. The ink composition can also be applied in a manner similar to the application of the outer primary coating on an ortical glass filter drawing and coating tower.
- [0101] Because of the versatility of the presently invented this costing composition, this composition is very well suited for use on costed optical glass fibers in ribbon assembles. Multiple individual strands of costed fiber are packaged into larger structures such as ribbons and cables to maximize efficiency. One skilled in the art will easily be able to use the disclosure provided hreein to prepare a novel ribbon assembly containing at its eaton eof the improved link costed octical glass fibers for the discrete applications. The novel ribbon assembly made according to this invention can be used in telecommunication systems. Such telecommunication systems typically include ribbon assemblies containing optical glass fibers, transmitters, receivers, and existince. The ribbon assembly containing the coaded optical gives fibers are
- tratecommunication systems. Such teacommunication systems typically include inborn assemblies containing optical gass fibers, frammitters, receivers, and switches. The fibbon assembly containing the casted optical gisss fibers are the fundemental connecting units of telecommunication systems. The ribbon assembly can be buried under ground or water for ong distance connections, such as between offers. The ribbon assembly can also be used to connect directly to residential homes.

 6 101021 The novel ribbon assembly made according to this invention can also be used in cable television systems.
- [9103] The ink compositions according to the present invention are surpsisingly suitable for formulating radiation-curable coloured coating compositions for forming protective and/or decorative coatings having a thickness greater than about 10 microns, and up to about 50 microns. While ink coatings on optical glass fizers usually have a thickness of
- about 3 to about 10 microns, the cure speed of the link compositions according to the present invention have been described above using a thickness of 20 microns. Thus, at the thinner thicknesse usually sected for forming an ink ceating on an optical glass fiber, such as from about 3 to about 10 microns, the MEK rub value will be significantly greater than the MEK rub value measured at a 20 micron thickness. In general, as the thickness of the ink coating is increased, the cure stread informasses.
- [0104] Thicker colored coatings tend to have a decreased concentration of pigment, based on the total weight of the dolored coating, compared to hit coating. Because the colored coating is applied in at a thickness greater than ink coating, a lower concentration of pigment is required to obtain the desired color.
 - [0105] The colored coating composition exhibits an enhanced cure speed and therefore is commercially feasible for use on high speed production lines, such as for appliances. The colored coating composition is also suitable for application to stroraft and promobile parts.
- 50 [0106] The invention will be further explained by way of the following examples, without being limited thereto.

Example !

[0107] A radiation-curable base composition was first formed by combining the components shown in Table 1.

Table 1

Component	Amount (% by weight of total composition)
Ethoxy Diacrylate Monomer ¹	42.85
Aliphatic Urethane Diacrylate Oligomer ²	25.25
Pentaerythritol Tetra-acrylate	12.55
1,6-Hexanediol Diacrylate	1.68
Isobornyl Acrylate	3.88
Phenoxy Ethyl Acrylate	3.88
Butylhydroxy Toluene	0.52
Benzophenore	8.33
2-Methyl-1-(4-(Methylthio)-phenyl)-2-(4-Morpholinyl)-1-Propanone	1.04
¹ CN120, (Sartomer)	
² Ebecryl 264, (Radcure)	

[0108] 96% of this base composition was then combined with the 4% of CGI 819 (Ciba Gelov) to form a radiation-curable composition. 88 wt.% of the radiation-curable composition was combined with 9 wt.% of blue, and 3 wt.% of white pigment to form a radiation-curable ink composition.

[0109] A 20 micron thick drawdown of the link compositions was formed on a mylar film and then exposed to 0.1 J/cm² UV light to form a sample film.

[0110] The following properties of the sample film was tested;

(1) MEK rub:

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- (2) Unreacted Coating: and
 - (3) Undercut.

[0111] The MEK rub test represents the degree of cross-linking that occurred in the sample. In general, the greater the number of rubs required to disintegrate the sample, the greater the degree of cross-linking, and the faster the ink composition cured.

- [0112] The unreacted coating test was a visible test for the presence of uncured coating on the sample. The presence of uncured coating signifies that a higher radiation dose may be required or that more of the cure speed enhancing photoinitiator is required to further increase the cure speed so that the link composition is suitably cured at the selected radiation dose.
 - [0113] The undercut test denotes whether the sample disintegrated after the MFK rub test
 - [0114] The test sample showed 200 MEK (rubs), no unreacted composition and no undercut.

Example II and III, and Comparative Example A

[0115] Two radiation-curable carrier systems were first formed by combining the components shown in Tables 2 and 3.

Amount (% by weight of total system
ranount (rob) noight or total by stein
33.9
19.9
9.96
3.06
3.06
13.64
1.14

Table continued

Component	Amount (% by weight of total system)
2,6-di-tert-Butyl-Methyl-Phenol	0.57
2-Methyl-1-(4-(Methylthio)phenyl)-2-(4-Morpholinyl)-1-Prapanone	4.55
Ink Cure Speed Enhancing Photoinitiator CGI 819	1.14
Acrylated Benzochenone	9.09

Table 9 Carrier System BB

Table 3 Carrier System	III DD
Component	Amount (% by weight of total system
Epoxy Acrylate monomer	35.93
Aliphatic Urethane Discrylate Oligomer	21.14
Pentaerythritol Tetra-acrylate	10.55
sobornyl Acrylate	3.24
Phenoxy Ethyl Acrylate	3.24
1,6-Hexanediol Discrylate	11.52
Silicone Discrylate	2.31
2,6-di-tert-Butyl-Methyl-Phenol	0.58
Phosphine Oxide, Di-Phenyl(2,4,6-Trimethy) Benzoyl)	11.52

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The radiation-curable carrier system AA was combined with the pigments shown in Table 4 to form radiation-curable ink compositions. The radiation-curable carrier system BB was combined with the pigments shown in Table 5 to form radiation-curable ink compositions. For comparison, commercially available ink coating compositions shown in Table 6 were utilized (comparison experiment A).

[0116] 20 micron thick drawdowns of the ink compositions were formed on mylar films and then exposed to 0.1 J/cm²
UV light to form sample films. The properties of the films was tested and the results are shown in Tables 4 through 6.

based on total weight of composition field state of the	Component (weight *	White Red	Red	Violet	Viciet Yellow Rose		Black	Slate	Black Slate Orange	Blue	Green	Brown	Agua
en (A) En (A)	based on total weight												
Se. 37 72.14 61.77 79.40 69.23 62.27 81.44 69.00 65.5 61.77	of composition)												
13.63 21.06 16.23 20.29 17.00 10.73 11.75 18.36 12.00 14.50 18.23	Radiation-Curable	86.37	78.14		79.80	83.00	89.39	88.27		96.00	85.5	91.79	85.35
13.63 31.86 16.23 20.39 17.99 10.71 11.73 12.36 12.00 14.50 19.21	Carrier System (A)												
Pack	Pigment	13.63	21.86	16.23	20.30	17.00	10.71	11.73	38.36	12.00	14,50	18.21	14.65
14.0 15.0	Test Results						L						
Natcle 100 2004 2004 2004 2004 110 2004	Viscosity (mPa.s)	2410	1820	2150	2140	2360	2300	2440	2260	2420	2160	2300	2265
MATCIX pass <	WEK (ruba)	200±	200+	200+	200+	200+		170	200+	130	200+	1002	\$00z
Watelk page 914ger page 914ger page								700		190			
Otter Přillinsty good good good good good good good goo	Release From Matrix	paes	pass	siight	pass	pasa	pags	basa	рава	page	pass	pass	page
Outst Primary good good good good good good good goo	Material			zipper									
75 74 77 63 84 83 61 60 61 79 79 70 78 75 77 75 75 75 75 75 75 75 75 75 75 75	Adhesion to outer primary	poof	рооб	2000	. pood	Bood	good	poof	poof	poob	good	pood	good
15 74 77 83 84 83 81 81 80 81 73 75 75 75 75 75 75 75 75 75 75 75 75 75	Coating				_		_						
73 67 73 79 79 79 77 78 75 77 75 75 Was 80	RAU In N, (*)	7.5	3.4	11	83	84	83	10	83	9.6	6.1	7.9	13
1	RAU In 0, (#)	73	6.3	73	7.5	96	75	7.	78	25	17	75	75
	Average RAU in N2 (%	Mas (200										

*

able 5

												-
Component (weight &	White	ped	Violat	Yellow	Rose	Black	Slate	Black Slate Orange Blue	Blue	Green Brown Aqua	Brown	Agus
based on total weight												
of composition)												
Radiation-Curable	85.01	96.24	82.14	97.78	81.30	86,22	67.10	79.91	86.80	84.05 79.97		83.14
Carrier System (A)												
Pigment	14.99	14.59 24.04 17.86	17.86	22.23	18.70	11.78 12.90 20.11	12.90		13.3	15,95 30.03	20.02	16.86
Test Results										- 1		
Viscosity (mPa.s)	2420	1580	2120	2250	2260	2280	2474	3300	2370	2260	2200	2320
MSK (rubs)	200+	100¢	1005	200+	300+	200+	2004	1005	700₹	200+	+005	+005
Release From Matrix	pass	Bsed	pass	page	gged	раяв	pass	pana	раза	paes	pass	page
Material												
Achesion to- outer primary	pood	poof	good	good	poof	poof	good	gond	pood	pood	900	9006
Coating												
RAU In N, (4)	8.1	81	97	58		88	83	87	63	98	88	83
RAU in air (*)	80	69	74	7.4	78	78	19	75	26	76	7.	7.9

Average RAU in N2 was 87; average RAU in air (%) was 79

able 6

11 11 11 11 11	White Bod	Bed	Violet	Violet / Yellow Rose	Rose	Black	Slate	Black Slate Orange	Blue	Green	Brown Agua	Agrua
Teac wearing	1											
MER (ruba)	200	200	155	108	200	180	155	145	133	300	185	135
Release From Matrix	poor	poor	poor	2000	poor	poor	poor	: boot:	1000	porr	poor	poor
Material												
Adhealon to outer primary	good	poof	good	poof	рооб	Book	poof , poof	poof	Bood	poof	goog	pood
Coatting												
RAU in N, (%)	69	T.	70	7.4	78	9	78	47	22	74	7.4	59
RAU in air (*)	54	89	74	73	22	7.8	17	11	5.1	75	E C	95

Average RAU in N_1 (%) was 74 Average RAU in air (%) was 70

[0117] By comparing the test results of Tables 4 and 5 with Table 6, it can be easily observed that the presence of a

suitable photorilitator provided a drematic increase in the average %FAU, when curing in the presence and absence of oxygen. It can also be observed that the presence of a suitable photorilitator provided the ink coeting with good adhesion to the outer primary coating in combination with providing good release from the matrix material even when cured at a low irradiation dose. Thus, the link coetings formed according to the present invention are suitable for use in a rebon assembly when breaks not the individual coosid optical glass fibers is deserted.

Example IV

[0118] An ink coating composition was prepared by mixing the following components:

Ebecryl 3700 (epoxy scrylate) 51.0		Component:	Percent by weight of total composition;
15 Irgacure 8191 1.0 1rgacure 907 3.0 Benzophenone 4.0 Blue pigment (Penn Color) 1.5 BYK 3332 0.5		Ebecryl 3700 (epoxy acrylate)	51.0
15 rgacure 907 3.0 Benzophenone 4.0 Blue pigment (Penn Color) 1.5 BYK 333 ² 0.5		alkoxylated aliphatic diacrylate	30.0
rgacure 907 3.0 Benzophenone 4.0 Blue pligment (Penn Color) 1.5 BYK 333 ² 0.5		rgacure 8191	1.0
Blue pigment (Penn Color) 1.5 BYK 333 ² 0.5	15	rgacure 907	3.0
BYK 333 ² 0.5		Benzophenone	4.0
1		Blue pigment (Penn Color)	1.5
20 1 Irgacure 819 is supplied by Ciba Gelgy		BYK 3332	0.5
	20	¹ rgacure 819 is supplied by Cit	oa Geigy

² BYK 333 is a polyether modified dimethyl polysiloxane.

Example v

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[0119] Another ink coating composition was prepared by mixing the following components:

Component:	Percent by weight of total composition:
polyether urethane acrylate: PTMG / TDI / HEA1	60.0
ethoxylated bisphenol-A diacrylate	25.0
N-vinyl caprolactam	25.0
trimethylo/propanetri-acrylate	4.0
Lucirin TPO ²	2.0
Benzophenone	3.5
Blue pigment (Penn Color)	1.5
silicone oli ³	0.5
1 reaction product of polytetramethylene glycol,	toluene diisocyanate and hydroxyethyl

acrylate

- 45 [0120] To test the lok compositions, glass plates were coated with a 75 micron thick, outer primary coating which was cured with a D-lang full denity. Next, the 75 micron thick drawdowns of the hick compositions, shown in examples IV and IV, were applied to coated glass plates, and then irradiated with ultraviolet light by a combination of a D largn with an energy of 1000 m Ucm² in the wavelength area up to 450 m and a H lamp with an energy of 500 m Ucm² in the wavelength area from 200 to 400 mm to moduse cured his films.
- 50 [0121] With these test plates, the adhesion of the ink to the outer primary coating was determined.
 - [0122] In order to test the adhesion (or break-out) of the matrix material from the links, coated and linked glass plates were used, and a matrix material was coated thereon. The matrix was cured with a D-lamp at 1 J/cm².
 - [0123] The energy output of the D and the H lamp were measured by a EIT UV-cure light bug.
- [0124] The curred links, described in the examples IV and V, showed a good adhesion to the outer primary coating, and the matrix showed a good break-out from the link layers.

² 2,4,6-trimethylbenzoyldiphenylphosphine oxide [Lucirin TPO by BASF]

³ silicone oil = Dow Corning 57/Dow Corning 190 mixture, whereby Dow Corning 57: polyether modified dimethyl polysiloxane, and Dow Corning 190: siliconeglycol copolymer

Test procedures

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[0125] The adhesion strenght of the cured ink coating and peelability of the matrix material were measured using the following methods:

- (1) Adhesion strenght
- [0126] The ink compositions, as were applied to an outer primary coating, were cut into with a knife in order to evaluate whether a good adhesion was obtained or not. A good adhesion was denoted as (+), a bad adhesion as (-); intermediate values were given ratings with (+) and (-).
 - (2) Peciability test
- [0127] The peelability of the matrix material from the ink composition was judged. The cured films were cut into with a knife in order to evaluate whether a good break-out was obtained or not. The peclability of the matrix material from the ink layer was evaluated by observation of the samples using the naked eye and touching the samples with the finger to examine the presence or absence of residues of the matrix on the ink layer. If neither observation confirmed any residues remaining on the ink coating when pulling away the matrix/ribbon material, the peelability was judged as good (+); if the presence of the matrix was confirmed or if the ink layer remained to the matrix, the peelability was judged as 20 bad (-).
 - [0128] The following test examples were studied and the results of the test procedures are given in Table 7:

Comparitive example B:

[0129] The composition of Example II without benzophenone.

Example V; as described above

В

Example VI: 30

[0130] The composition of Example V + 3% Ebecryl 350 wherein Ebecryl 350 (UCB-Radgure) is an acrylated silicone (difunctional).

Example VII

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[0131] The composition, which can be used as a colored outer primary coating, was prepared by mixing the following naredients.

	Component:	Percent by weight of total composition:
40	polyester based urethane acrylate	22
	polyether based urethane acrylate	20
	isobornyl acrylate	14
	ethoxylated TMPTA	17
45	ethoxylated bisphenol A diacrylate	16
10	Irgacure 907	3
	Irgacure 819	1
	Blue pigment (Penn Color)	1
	Titanium dioxide	2
50	1,6-nexamethylene discrylate	3
	benzophenone	4

Table 7

Release (of the matrix from the lnk) Adhesion (of the ink to the outer primary coating)

Table continued

		Release (of the matrix from the lnk)	Adhesion (of the ink to the outer primary coating)
	٧	++	4.
6	VI	++	+
2	100		

[0132] The use of benzophenone alone results in a good break-out. Eberory 350 does not improve this substantially, but nevertheless, docreases the schesion of the link conting to the outer primary coating, in other words; it is greferred not to use considerable amounts of release agents (such as Ebercyl 350) as this reduces the adhesion strength of the intk coating on the outer primary coating.

[0133] Furthermore, these experiments show that benzophenone substantially improves the break-out of the matrix material from different his types.

15 Example VIII

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[0134] "If micron thick drawdowns of two commercially available radiation-curship outer primary ceating compositions were cured on Mylar sheats by exposure to 1.0 Jcm² Ur radiation to form outer primary films. Samples were made by cutting the cured time two 2 inch by 2 inch squares. The samples were then placed in a desiccator for a minimum of 15 minutes. The initial weight of each of the samples were measured. 3 samples were submerged into each of eight different monomers and weighted at intervals of 1 seconds, 5 seconds, 10 seconds and 30 seconds (total elapsed time). The percent weight change of each of the samples and the average weight change of the three samples for each monomer was acculated and the results are shown in Table 3.

Table 8

	Commercial Outer Primary Coating No.	Outer Pri	mary Coati	ng No. 1	Commercial	Outer Pri	Commercial Outer Primary Coating No.	19 No. 2
	Percent Change In Weight After 30	inge In We	ight After	30	Percent Ch	ange in We	Percent Change in Weight After	30
	Seconds				Seconds			
Monomer	Samp. 1	Samp. 2	Samp. 3	Average	Samp. 1	Samp. 2	Samp. 3	Average
HDDA	+.58	+.5B	+.39	+.52	+.52	+.58	+.39	+.5
BPAEDA	4.8	+1.13	+1.14	+1.02	+.46	+.37	+.33	+.39
SA1002	+1.03	+.46	+.64	+.71	+.98	+.26	+.75	+.66
IBOA	+.25	+.37	+.26	+.29	+.43	+.25	+.11	+.26
SR504A	+1.14	1.87	+.19	+.73	+.14	+.48	+.27	+.3
PEA	4.8	+.88	+.93	+.87	+.85	+.56	+.37	+.59
n-Vinyl	+2.32	+2.91	+3.11	+2.78	+2.71	+2.72	+3.21	+2.88
Formamide								
IDA	+.23	+.25	+.30	+.30	+.14	+.13	+.33	+.19
	Percent Change in Weight After 10	ange in We	ight After	10	Percent C	ange in We	Percent Change in Weight After	10
	Seconds				Seconds			
Monomer	Samp. 1	Samp. 2	Samp. 3	Average	Samp. 1	Samp. 2	Samp. 3	Average
HDDA	+.55	+.75	+.22	+.51	+.72	+.38	55.4	+.55
BPAEDA	+1.24	+:65	+,58	+.82	+.33	+.44	+.3	+.36
_	_	_		_	-			

SA1002	+.53	+.46	+.38	+.46	+.29	+.33	+.82	+.48
IBOA	+.31	+.35	+.29	+.32	+.24	+.03	4.5	+.26
SR504A	+.76	+.93	+.31	+.53	+.09	+.42	+.27	+.26
PEA	+.74	+.38	+.55	+.66	+.75	+.32	+.42	٠.5
n-Vinyl	+1.69	+2.31	+2.54	+2.18	+1.97	+1.96	+2.62	+2.18
Formamide								
IDA	+.26	+.28	+.30	+.28	+.14	+.08	+.18	+.13
Company of the Control of the Contro	Percent Change in Weight After	ange in We	sight After	5 .	Percent C	hange in We	Percent Change in Weight After	S
	Seconds				Seconds			
Monomer	Samp. 1	Samp. 2	Samp. 3	Average	Samp. 1	Samp. 2	Samp. 3	Average
нора	+.78	+.54	+.27	+.53	+.8	+.61	+.71	+.71
BPAEDA	+.93	+1.07	+.74	+.91	+.26	+.37	+.33	+.32
SA1002	+.87	+.46	+.29	+.54	+.53	+.51	+.58	+.54
IBOA	+.2	+.32	+.19	+.24	+.34	+.34	+.22	+.3
SR504A	+1.83	+.76	+.31	+.97	+.26	+.28	+.27	+.27
PEA	+.67	+.41	+.24	+.44	+.5	+.6	+.48	+.53
n-Vinyl	+1.13	+1.92	+1.69	+1.58	+1.44	+1.34	+2.02	+1.6
Formamide								
IDA	+.2	+114	+.15	+.16	+.21	+.14	+.22	+,19

	Percent Cha	inge in We	ight After	1 Second	Percent Ch	nange in We	Percent Change in Weight After 1 Second Percent Change in Weight After 1 Second	1 Second
HDDA	+.45		+.17	+.34	+.88	+.46	+.39	+.58
BPAEDA	+.65	+1,1	+1.07	+.94	+.26	+.3	+.33	+.3
SA1002	+.87	+.62	+.44	+.94	+.37	+.29	+.58	+.41
IBOA	+.28	+.4	+.42	+.37	+.3	+.28	+.11	+.23
SR504A	+1.03	+.64	+.31	+.66	+.26	+.11	+.17	+.18
PEA	+.43	+.24	+.38	+.35	+.46	+.35	+.34	+.38
n-Vinyl	4.59	+.73	+.98	+.77	+.49	+.59	+1.05	+.71
Formamide								
IDA	+.2	+.25	+.11	+.19	+.14	+.14	+. 15	+. 11
			1					

[0135] The test results in Table 8 demonstrate that the monomers present in radiation-curable ink compositions exhibit different levels or desorption into different course ordality available outper primary costilings. In general, the greater the association into the outer primary coating, the greater the resulting adhesion between the cured ink coating and the outer primary coating.

Test Procedures

%RAU

- © [0136] This is an FTIR method for determining the degree of cure on the surface of a coating to a depth of 1 to 3 µm, and following exposure to a prodetermined dose of UV radiation. A drawdown of the sample material is first formed on a glass plate. The drawdown is the nexposed to the predetermined dose of UV radiation to cure the drawdown and form a coating. The coating is cut into strong, permissely 10 mm by 50 mm.
- [0137] An Infrarod spectrum of the unoured, fluid sample and an infrared spectrum of the cured sample are obtained.

 Infrared spectrum well know and any infrared spectrum of the unoured liquid sample is measured. For most acrylate based obtains the infrared spectrum.

 [0138] The net peak erea of the acrylate unsaturation absorbance for the unoured liquid sample is measured. For most acrylate based obtains, the absorbance at about 810 cm⁻¹ and should be used. However, if the obtains contain outside as slovate or other component which absorbs strongly at or near 810 cm⁻¹, an alternative acrylate absorbance peak can be used. The absorbances at about 1410 cm⁻¹ and about 1458 cm⁻¹ have been found to be satisfactory. The not peak of an according to the specific peak of the substitution of the contribution of the
 - [0139] A reference area is then determined. The reference absorbance should not change in intensity as the figuld sample is cured. Many formulations have an absorbance in the range of about 750 to about 750 to com³ that can be used as a reference absorbance. The net peak area of the reference absorbance is measured.
- 25 [0140] The ratio of the acrylate absorbance to the reference absorbance for the uncured, liquid sample is determined using the following formula:

$$R_r = A_{rr} / A_{pr}$$

where

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AAL is the net peak area of the acrylate absorbance,

App is the net peak area of the reference absorbance, and

R_L is the area ratio for the liquid sample.

[0141] The ratio of the acrylate absorbance to the reference absorbance for the cured sample is determined using the following formula:

$$R_c = A_{ac} / A_{RC}$$

where

6 A_{4.0} is the net peak area of the acrylate absorbance.

ARC is the net peak area of the reference absorbance, and

Ro is the area ratio for the cured sample.

[0142] The degree of cure as a percent reacted acrylate unsaturation (%RAU) is determined using the following formula:

$$RAU = [(R_L - R_C) \times 100\%] / R_L$$

MEK rub Test

[0143] The test procedures described in ASTM D 4752 was followed with the following exception. Light finger pressure was used on the material being rubbed against the test sample.

[0144] While the claimed invention has been described in detail and with reference to specific embodiments thereof,

It will be apparent to one of ordinary skill in the art that various changes and modifications can be made to the claimed invention without departing from the scope thereof.

5 Claims

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- 1. A radiation-curable ink coating composition edapted to provide a color coding to a coated optical glass fiber comprising
- a) an oligomer having at least one radiation curable group
- b) a reactive diluent
 - c) at least about 1 wt.% of homolytic photoinitiator for radical polymerisation

 - e) additives
- wherein the composition comprises more than 2 weight% of a benzophenone-type compound, wherein said weight% is with respect to the total weight of said radiation-curable ink composition.
 - 2. A composition according to claim 1, wherein more than 3 weight% of a benzophenone-type compound is present,
- 3. A composition according to any of claim 1 or 2, wherein the benzophenone-type compound is an organic compound with a molecular weight between 170 and 500, consisting essentially of benzophenone or a subtituted benzophenone group.
- A radiation-curable ink coating composition according to any of claims 1-3, wherein the benzophenone-type compound is not a ring-bonded benzophenone.
 - 5. A composition according to any one of claims 1-4, wherein less than 0.1 weight% of an amine is present,
 - 6. A composition according to any one of claims 1-5, wherein the homolytic photoinitiator is represented by formula (1),

wherein:

Ar represents a carbon containing compound having at jeast one gromatic functional group;

P is bonded directly to an aromatic functional group present in Ar3;

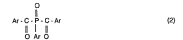
C1 is bonded directly to an aromatic functional group present in Ar1; and

C2 is bonded directly to an aromatic functional group present in Ar2, and

wherein at least one of the carbon containing compounds Ar has a molecular weight and molecular structure that is capable of forming at least one of the following free-radicals upon exposure to ultraviolet radiation:

or

7. A composition according to claim 6, wherein the homolytic photoinitiator is represented by formula (2),



wherein Ar is a phenyl group, optionally substituted with one to three alkyl groups having 1-4 carbon atoms.

- 8. A composition according to any one of claims 1-7, wherein compound e) comprises at least a jubricant.
- A radiation-curable lnk composition according to any one of claims 1-8, adapted for coating optical glass fibers and being formulated from a composition comprising:
 - from about 20 to about 70 weight % of at least one radiation-curable oligomer:
 - from about 5 to about 80 weight % of at least one radiation-curable diluent;
 - from about 4 to about 20 weight % of a homolytic photoinitiator;
- from about 1 to about 20 weight % of a pigment, and more than about 2 weight % of a benzophenone-type compound.
 - 10. A method for curing an ink composition according to any one of claims 1-9 on a coated optical fiber, wherein the composition is cured with a combination of a D and a H lamp.
- 25 11. A coated and colored optical glass fiber, comprising:
 - an optical class fiber:

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- an inner primary coating adjacent to the surface of said optical glass fiber:
- an outer primary coating adjacent to the surface of said inner primary coating; and
 - an ink coating adjacent to the surface of said outer primary coating, wherein the ink coating is the cured ink coating composition, according to any one of claims 1-9.
- 12. A coated and colored optical glass fiber, comprising:
 - an optical glass fiber;
 - an inner primary coating adjacent to the surface of said optical glass fiber,
 - a colored cuter primary coating adjacent to the surface of said inner primary coating wherein the outer primary coating is the cured ink coating composition according to any one of claims 1-9.
- 0 13. A glass fiber according to anyone of claims 11-12, wherein the cursed ink costing composition comprises a persent reacted acrylate unsaturation of at least 70% when the composition is cured by exposure to a radiation dose of 0.125 joules aquare centimeter.
- 14. A glass fiber according to claim 13, wherein the cured ink coating composition comprises a percent reacted acrylate unsaturation of at least 75%.
 - 15. A ribbon assembly comprising:
 - a plurality of coated optical glass fibers; and
 - a matrix material which binds said plurality of coated optical glass fibers together, wherein at least one of said coated optical glass fiber according to anyone of claims 11-14.

Revendications

 Composition de revêtement d'encre durcissable par irradiation, destinée à donner un codage coloré à une fibre de verre optique enrobée, comprenant

- a) un oligomère comportant au moins un groupe durcissable par irradiation.
- h) un diluant réactif
- c) au moins environ 1 % en poids d'un photoamorceur homolytique pour polymérisation radicalaire
- d) un pigment, et
- 5 e) des additifs

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dans laquelle la composition comprend plus de 2 % en polds d'un composé de type benzophènone, ledit pourcentage en poids étant par rapport au poids total de ladite composition d'encre durcissable par irradiction.

- Composition selon la revendication 1, dans laquelle sont présents plus de 3 % en poids d'un composé de type benzophénone,
 - Composition selon la revendication 1 ou 2, dans laquelle le composé de type benzophénone est un composé
 organique ayant une masse moléculaire comprise entre 170 et 500, et consistant essentiellement en benzophénone
 ou en un crouse benzophénone substitué.
 - Composition de revêtement d'encre durcissable par irradiation selon l'une quelconque des revendications 1 à 3, dans laquelle le composé de type benzophénone n'est pas une benzophénone liée à un cycle.
- Composition seion l'une quelconque des revendications 1 à 4, dans laquelle est présent moins de 0,1 % en poids d'une amine.
 - Composition seion l'une quelconque des revendications 1 à 5, dans laquelle le photoamorceur homolytique est représenté par la formule (i)

dans laquelle :

- Ar représente un composé carboné comportant au moins un groupe fonctionnel aromatique; P est lié directament à un groupe fonctionnel aromatique présent dans Ar³; et C² est lié directament à un groupe fonctionnel aromatique présent dans Ar¹; et C² est lié directement à un groupe fonctionnel aromatique présent dans Ar², et
- 40 dans laquelle au moins l'un des composés carbonés Ar a une masse moléculaire et une structure moléculaire capables, lors d'exposition à un rayonnement ultraviolet, de former au moins l'un des radicaux libres suivants :

Ar1-C10+:

45 Ar2-C2O :

ou

Ar3.PO+

7. Composition selon la revendication 6, dans laquelle le photoamorcaur homolytique est représenté par la formule (2).

- dans laquelle Ar est un groupe phényle éventuellement substitué par un à trois groupes alkyle ayant de 1 à 4 atomes de carbone.
 - Composition selon l'une quelconque des revendications 1 à 7, dans laquelle le composé e) comprend au moins un lubrifiant.
 - Composition de revêtement d'encre duroissable par irradiation selon l'une que conque des revendications 1 à 8, destinée au revêtement de fibres de verre optiques et étant formulée à partir d'une composition comprenant :
 - d'environ 20 à environ 70 % en poids d'au moins un oligomère durcissable par irradiation ;
 - d'environ 5 à environ 80 % en poids d'au moins un dituant durcissable par irradiation ; d'environ 4 à environ 20 % en poids d'un photoamorceur homolytique ;
 - d'environ 1 à environ 20 % en poids d'un pigment ; et
 - plus d'environ 2 % en poids d'un composé de type benzophénone.
- Procédé pour le durcissement d'une composition d'encre selon l'une quelconque des revendications 1 à 9 sur une fibre optique revêtue, dans lequel la composition est durcie avec une combinaison d'une lampe D et d'une lampe H,
 - 11. Fibre de verre optique revêtue et colorée, comprenant :
 - une fibre de verre optique :

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- un revêtement primaire interne adjacent à la surface de ladite fibre de verre optique ;
- un revêtement primaire externe adjacent à la surface dudit revêtement primaire interne ; et
- un revêtement d'encre adjacent à la surface dudit revêtement primaire externe, le revêtement d'encre étant la composition de revêtement d'encre durcie selon l'une quelconque des revendications 1 à 9.
- 12. Fibre de verre optique revêtue et colorée, comprenant :
 - une fibre de verre optique ;
 - un revêtement primaire interne adjacent à la surface de ladite fibre de verre optique ;
 - un revêtement primaire externe coloré adjacent à la surface dudit revêtement primaire interne, le revêtement primaire externe étant la composition de revêtement d'encre durcle seion l'une quelconque des revendications 1 à 9.
- 13. Fibre deverte selon ia revendication 11 ou 12, dans laquelle la composition de revêtement d'encre durcle comprend un pour certage d'insaturation d'acrylate ayant réagi d'au moins 70 %, quand la composition est durcle per exposition à une dose d'irradiation de 0.125 joule par cm².
 - 14. Fibre de verre selon la revendication 13, dans laquelle la composition de revêtement d'encre durcle comprend un pourcentage d'insaturation d'acrylate ayant réagl d'au moins 75 %.
 - 15. Ensemble sous forme de ruban comprenant :

plusieurs fibres de verte optiques revêtues : et

une matière de matrice qui lie entre elles lesdites plusieurs fibres de verre optiques revêtues,

dans lequel au moins l'une desdites fibres de verre optiques revêtues comprend une fibre de verre optique revêtue et colorée selon l'une quelconque des revendications 11 à 14,

Patentansprüche

- Durch Strahlung h\u00e4rbare Tintenbeschichtungszusammensetzung, die eingerichtet ist, eine beschichtete optische Glasfaser mit einer Farbkodierung zu versehen, umfassend
 - a) ein Oligomer mit mindestens einer durch Strahlung härtbaren Gruppe,
 - b) ein reaktives Verdünnungsmittel,
 - c) mindestens etwa 1 Gew.-% eines homolytischen Photoinitiators zur radikalischen Polymerisation,
- d) ein Pigment und

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e) Additive,

wobei die Zusammensetzung mehr als 2 Gew.-% einer Verbindung vom Banzophenon-Typ umfaßt, wobei die Gew.-Prozente bezüglich des Gesamtgewichts der durch Strahlung härtbaren Tintenzusammensetzung sind.

- Zusammensetzung gemäß Anspruch 1, wobei mehr als 3 Gew.-% einer Verbindung vom Benzophenon-Typ vorliegt.
 - Zusammensetzung nach einem der Ansprüche 1 oder 2, wobei die Verbindung vom Benzophenon-Typ eine organische Verbindung mit einem Molekulargewicht von zwischen 170 und 500 ist, bestehend im wesentlichen aus Benzophenon oder einer substitutierte Benzophenon-Gruppen.
 - Durch Strahlung h\u00e4rtbare Tintenbeschichtungszusammensetzung gem\u00e4\u00df einem der Anspr\u00fcche 1 bis 3, wobei die Verbindung vom Benzophenon-Typ nicht ein Ring-gebundenes Benzophenon ist.
 - Zusammensetzung gemäß einem der Ansprüche 1 bis 4, wobei weniger als 0,1 Gew.% eines Amins vorliegt.
 - Zusammensetzung gemäß einem der Ansprüche 1 bis 5, wobei der homolytische Photoinitiator durch Formel (1) dargestellt ist

worln

Ar eine Kohlenstoff-enthaltende Verbindung mit mindestens einer aromatischen funktionalen Gruppe darstellt, P direkt an eine aromatische funktionale Gruppe, vorliegend in Ar3, gebunden ist,

C1 direkt an eine aromatische funktionale Gruppe, vorliegend in Art, gebunden ist und

C2 direkt an eine aromatische funktioneile Gruppe, vorliegend in Ar2, gebunden ist, und wobei mindestens eine der Kohlenstoff-entheitenden Verbindungen Ar ein Molekultargewicht und Molekültsruktur mit der Befähigung aufwelst, mindestens eines der folgenden freien Radikale nach Aussetzung ultravloietter Strahlung zu bilden,

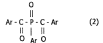
45 Ar1-C1O•

Ar2-C2O+

oder

Ar3-PO+

7. Zusammensetzung gemäß Anspruch 6, wobei der homolytische Photoinitiator durch Formel (2) dargestellt ist



- worin Ar eine Phenylgruppe, gegebenenfalls substituiert mit einem bis drei Alkylresten mit 1-4 Kohlenstoffatomen, ist.
 - 8. Zusammensetzung gemäß einem der Ansprüche 1 bis 7, wobei Verbindung e) mindestens ein Schmiermittel umfaßt,
 - Durch Strahlung h\u00e4ribare Tintenzusammensetzung gem\u00e4\u00df einem der Anspr\u00fcche 1 bis 8, eingerichtet zum Beschichten von optischen Glasfasern und aus einer Zusammensetzung formuliert, umfassend:

von etwa 20 bis etwa 70 Gew. % von mindestens einem durch Strahlung härtbaren Oligomer, von etwa 5 bis etwa 80 Gew.% von mindestens einem durch Strahlung härtbaren Verdünnungsmittel, von etwa 4 bis etwa 20 Gew. % elnes homolytischen Photoinitiatiors,

20 von etwa 1 bis etwa 20 Gew.-% eines Pigments, und

- mehr als etwa 2 Gew.% einer Verbindung vom Benzophenon-Typ.
- Verfahren zum H\u00e4rten einer Tintenzusammensetzung gem\u00e4\u00df elnem der Anspr\u00fcche 1 bis 9 auf einer beschichteten
 optlischen Faser, wobel die Zusammensetzung mit einer Kombination von einer D- und einer H-Lampe geh\u00e4rtet wird.
- Beschichtete und gef\u00e4rbte optische Glasfaser, umfassend
 - eine optische Glasfaser,
 - eine innere Primärbeschichtung angrenzend an die Oberfläche der optischen Glasfaser,
 - eine äußere Primätbeschichtung angrenzend an die Oberfläche der inneren Primätbeschichtung und eine Tintenbeschichtung angrenzend an die Oberfläche der äußeren Primätbeschichtung, webei die Tintenbeschichtung die gehärzte Tintenbeschichtungszusammensetzung gemäß einem der Ansprüche 1 sie 9 ist.
 - 12. Beschichtete und gefärbte optische Glasfaser, umfassend
 - eine optische Glasfaser,
- eine innere Primärbeschichtung angrenzend an die Oberfläche der optischen Glasfaser,
 - eine gefärche Bußere Primärbeschichtung angrenzend an die Oberfläche der inneren Primärbeschichtung, wobei die Bußere Primärbeschichtung die gehärtete Tintenbeschichtungszusammensetzung gemäß einem der Ansprüche 1 bis 9 ist.
- 10 13. Clasfaser gemäß einem der Ansprüche 11 oder 12, wobel die gehärtete Tirlenbeschichtungszusammensetzung einen Prozentanteil en regeiner Angralt-Ungesättighet von mindisstens 70% umfalt wenn die Tirlenbeschichtungszusammensetzung gehärtet ist nech aussetzung zu einer Härtungdosis von 0.125 Joules pro Qaudretzentimeter.
- 45 14. Glasfaser gemäß Anspruch 13, wobei die gehärtete Tirttenbeschichtungszusammensetzung einen Prozentanteil an reagierter Acrylat-Ungesättigtheit von mindestens 75% umfaßt.
 - 15. Bandeinheit, umfassend:
 - eine Vielzahl beschichteter optischer Glasfasern und
 - ein Matrixmaterial, welches die Vielzahl beschichteter optischer Glasfasern miteinander verbindet, wobei mindestens eine der beschichteten optischen Glasfasern eine beschichtete und gelärbte ontische Glasfaser gemäß Anspruch 10 bis 41 unrfaß!

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